SUMMARY OF THE QUALITY SYSTEMS COMMITTEE TELECONFERENCE SEPTEMBER 17, 1999

The Quality Systems (QS) Committee of the National Environmental Laboratory Accreditation Conference (NELAC) met by teleconference on September 17, 1999, at 1:00 p.m. Eastern Daylight Time (EDT). The meeting was led by its chair, Mr. Joe Slayton of EPA Region III. A list of action items is given in Attachment A. A list of participants is given in Attachment B. The list of parking lot issues includes 5 items following this meeting (Attachment C). Attachment D is a listing of frequently asked questions. Attachment E presents the QS Committee approach to handling comments, comment acknowledgment form letter, guiding principles for reviewing comments and the standard, and commenter template. Attachment F contains the QS Committee's response to comments addressed during this meeting. Attachment G is the NELAC Policy on Revisions of Standards. Attachment H is a proposed format for receiving comments on Chapter 5. Changes to the language in Chapter 5 proposed at this teleconference are reflected in version 5.10.13 of the standard. However, to avoid confusion within NELAC, since version 5.10.7 is the version provided for NELAC V voting, 5.10.13 is only being circulated within the QS Committee at this time (not attached to these minutes and will not be posted on the web). The purpose of the meeting was to discuss the review of comments received on Chapter 5 and to propose appropriate revisions in preparation for NELAC 5i.

DEPARTMENT OF DEFENSE

Mr. Glowacki is reviewing the DoD comments and is attempting to set up a meeting between QS Committee members and DoD representatives discuss the comments. It is important that these comments be resolved before the November 5 deadline.

MICROBIOLOGY

Ms. Bruch has reviewed the VWEA comments and will fax her comments in to Mr. Slayton. Some comments suggested the Microbiology section need to be simplified. QS Committee members feel the document has been simplified as much as possible but have concern that the laboratories may have some difficulty in understanding the language of the section. Dr. Siegelman will have Margo Hunt of EPA review the section for clarity and simplicity and clarity. Mr. Siders will also have the Illinois EPA review the document and Mr. Slayton will ask Ms. Darlene Raiford the review the section as well. Ms. Labie reminded the committee that the section on microbiology was adopted last year and is subject to the NELAC Policy on Revision of Standards (Attachment G) which discourages changes unless essential The committee also discussed the possibility of establishing a microbiology subcommittee for addressing these issues but decided that such a group is unnecessary at this time. The need for running positive QA controls on filters used for microbiological tests was emphasized by the committee members as a means of demonstrating that there is no agent on the filters which

might inhibit growth of cultures. The committee agrees that failing to run positive controls may lead to serious false negatives.

COMPARISON OF NELAC 5 VERSION OF CHAPTER 5 AND CURRENT VERSION

Dr. Siegelman has supplied the committee with a comparison of the WordPerfect versions of Chapter 5 used at NELAC 5 and the current 5.10.13 version being discussed by the committee. In this comparison the material that has been deleted since the NELAC 5 version appears in blue and the new material appears in red. Mr. Slayton urged all committee members to review this comparison document before the next teleconference on Wednesday, September 22. Any items which will need to be proposed for amendment at NELAC 5i should be identified.

GLOSSARY

Mr. Slayton has contacted Chairs of the other committees and the Chair of the NELAC Board of Directors and recommend that (1) that all items in the Glossary will be cross referenced to the specific committee concerned, and (2) when changes are proposed to entries in the Glossary, the committees which are affected by these changes will be notified and allowed to review the proposed change. The Board has not met to discuss this issue but initial response from several members has been favorable. Mr. Slayton will attend the next Board meeting on September 23 to discuss this issue.

COMMENT FORM (ATTACHMENTS E & F)

A new format has been proposed for submitting comments to the committee. Several public comments have stated the current table format is difficult to work with in some word processing software. A format using cells with headings has been proposed as a remedy. A draft format has been developed and is found in Attachment H. Ms. Labie advised the committee that NELAC is considering adopting the QS format for all comments on NELAC chapters. Changes to that format should be completed before the Board meeting on September 23.

ADDITIONAL COMMENTS

A review of terms in Chapter 5 which need to be reviewed for clarity and consistency were discussed. These items will be added to the "parking lot" and included:

"Such as" should have "but not limited to..." added where appropriate

There is confusion over the terms "independent standard," "alternate source," and "second source."

The sections on "Analyst Training" and "Verification" (5.6.2) are misplaced. Information needs to be combined into one section.

The committee reaffirmed its commitment to keeping Chapter 5 consistent with ISO 17025 when it is formally adopted by ISO.

Demonstration of capability for the laboratory and the work cell needs clarification.

Clarification between "sample custody" and "sample tracking" needs to be made.

The sections on Radiochemistry and Air Testing need to be finalized.

FUTURE MEETINGS

In addition to the telconference schedule given below, the committee discussed the need for a face-to-face meeting either immediately before or after NELAC 5i. After some discussion, the committee decided to meet after NELAC 5i to allow issues brought before the committee at NELAC 5i to be addressed.

QS TELECONFERENCE SCHEDULE

The following QS Committee teleconferences are scheduled before NELAC 5i:

	TIME(S)
DATE(S)	EASTERN
Sept. 22	2:30p-4:30p
Oct. 1	1p-3p
Oct. 15	1p-3p
Oct. 27	1p-3p
Nov. 3	1p-3p
Nov. 10	1p-3p

Telephone numbers for these teleconferences are included in an e-mail message from Mr. Slayton to the committee members on Friday, September 17, 1999 12:40 PM. The deadline for submitting committee materials for discussion at NELAC 5i is November 5. All QS Committee "homework" must be completed before that date in order to have comments considered at the NELAC 5i meeting in December.

ACTION ITEMS Quality Systems Committee September 17, 1999

Item No.	Action Item	Date to be Completed
1.	All committee members will review the comparison of Chapter 5 from NELAC 5 and the 5.10.13 version before the 9/22/99 teleconference.	Sept. 21, 1999
2.	Mr. Slayton will confer with Mr. Glowacki on review of the DoD comments.	Sept. 20, 1999
3.	Ms. Bruch to fax comments on microbiology section from VWEA and PA DEP to Mr. Slayton	Sept. 20, 1999
4.	Messers. Glowacki, Frederici, and Mendenhall will send comments on Wisconsin DNR to Mr, Slayton	Sept. 20, 1999
5.	Mr. Slayton will contact Mr. Porterfield for response to comments from US Navy.	Sept.20, 1999
6.	Committee members will contact reviewers for section on Microbiology: Dr. Siegelman will ask Margo Hunt, Mr. Siders will ask IL EPA, and Mr. Slayton will ask Darlene Raiford of VA to review section.	Sept. 20, 1999
7.	Improved format for receiving comments on Chapter 5 will be developed by committee (Ms Labie and Mr. Slayton).	Sept 20, 1999
8.	Mr. Slayton will update the comment table (Attachment F).	Sept. 22, 1999
9.	Mr. Beard to prepare draft minutes of the teleconference.	Sept. 20, 1999
10.	The next teleconference is September 22 from 2:30 p.m. to 4:30 p.m. EDT. The telephone number has been distributed by e-mail of 9/17/99 from Mr. Slayton.	Sept. 22, 1999

Attachment B

PARTICIPANTS Quality Systems Committee September 17, 1999

Name	Affiliation	Phone Numbers
Mr. Joe Slayton	USEPA, Region III, OASQA	T: 410-305-2653 F: 410-305-2698 E: slayton.joe@epamail.epa.gov
Ms. Mary K. Bruch	Mary Bruch Micro Reg. Inc.	T: 540-338-2219 F: 540-338-6785 T: Daughter - 301-469-7222 E: mkesterm@aol.com
Dr. Peter Delisle	Coastal Bioanalysts	T: 804-694-8285 F: 804-695-1129 E: pdelisle@coastalbio.com
Mr. Raymond J. Frederici	Severn Trent Laboratories	T: 708-534-5200 F: 708-534-5211 E: frederir@stl-inc.com
Mr. Clifford R. Glowacki (Absent)	Covenant	T: 916-643-0447 F: 916-643-0190 E: covenantea@stl-inc.com E: cglowacki@cerp-aiger.org
Dr. George Kulasingam	California Department of Health — ELAB	T: 510-540-2800 F: 510-849-5106 E: gkulasin@dhs.ca.gov
Ms. Sylvia S. Labie	Florida Department of Environmental Protection	T: 850-488-2796 F: 850-922-4614 E: labie_s@dep.state.fl.us
Mr. David Mendenhall	Utah Department of Health	T: 801-584-8470 F: 801-584-8501 E: dmendenh@doh.state.ut.us
Mr. Jeff Nielsen	City of Tallahassee Water Quality Division	T: 850-891-1232 F: 850-891-1062 E: nielsenj@mail.ci.tlh.fl.us
Mr. Scott D. Siders	Illinois Environmental Protection Agency	T: 217-785-5163 F: 217-524-0944 E: epa6113@epa.state.il.us
Dr. Fred Siegelman	US EPA, QAD	T: 202-564-5173 F: 202-565-2441 E: siegelman.frederic@epamail.epa.gov
Mr. Mike Beard (Contractor Support)	Research Triangle Institute	T: 919-541-6489 F: 919-541-7386 E: mebeard@rti.org

PARKING LOT ITEMS/ISSUES

Quality Systems Committee September 17, 1999

Items/issues will remain in the Parking Lot until they are completed.

The following items were added to the parking lot following the September 17, 1999 meeting:

- 1. Review terms in Chapter 5 for terms needing clarification, viz. "such as," "independent standard," "alternate source," "second" or "alternate source."
- 2. Combine "Analyst Training" and "Verification" into same section.
- 3. Clarify "Custody" versus "Sample Tracking."
- 4. Finalize Radiochemistry section.
- 5. Come to closure on Air Testing issues.

FREQUENTLY ASKED QUESTIONS Quality Systems Committee

September 17, 1999

Some Frequently Asked Questions Concerning NELAC QS (Chapter 5)

1. Question: If a mandated method (required by EPA or State Authority) is less stringent than the QS standards what do I follow?

Answer: The most restrictive/demanding.

2. Question: Do the QS standards require the use of any specific method?

Answer: No. QS does not require the use of a specific method/s. Chapter 5 allows the user to select an appropriate method. However, regulatory agencies may mandate the use of a specific method (See also Question 3).

3. Question: Do the QS standards allow for the use of the PBMS approach?

Answer: Yes. However, the QS standards may include additional QS checks/requirements (considered by NELAC to be essential) than those associated with a PBMS method for a given project. Such additional requirements would also apply to conventional or non-PBMS methods as well.

4. Question: Do the QS standards apply to small laboratories?

Answer: Yes. The standards include essential QC procedures and are applicable to environmental laboratories regardless of size and complexity. It is suggested that the amount of effort that will be required to attain the standards will be dependent on whether the laboratory already is operating under a quality system (with established and documented SOPs and QC procedures) more then upon the size of the laboratory.

5. Question: If my laboratory is measuring high level concentrations and is set-up (perhaps even optimized) to analyze at such levels and is only interested in whether a high level regulatory limit is exceeded, why do I have to determine a detection limit?

Answer: A detection limit is considered essential to verify (confirm and document) that the laboratory is actually able to detect and measure at the regulatory or decision limit. Detection limit determinations are also considered an important consideration with regard to the quantitation range selection particularly with regard to the choice of the concentration of the lowest calibration standard. Changes to the standard will be proposed at the January 1999

Interim Meeting, which no longer specify that the MDL (40 CFR Part 136) procedure be employed, unless it is mandated by the test method or applicable regulation. In the proposed revision, the term "detection limit" may not be the lowest concentration level attainable by a given analytical method, but rather that it is a concentration that is actually measurable (and verified) using the procedures, e.g., equipment, analytical method, routinely employed for sample analyses (could be relatively high concentration). The detection level should be appropriate or relevant for the intended use of the data. In some cases this will of necessity be the lowest concentration level attainable, e.g., low level drinking water or wastewater permit limits.

6. Question: Why are we revisiting the calibration and detection parts of the standards?

Answer: At NELAC IV the Quality Systems Committee received numerous comments that the calibration and detection parts of the standards were too prescriptive and were not consistent with a PBMS environment. The Committee has attempted to propose changes to the calibration and

detection parts of the standards that provide essential elements for those two quality system standards and that will support the anticipated needs of PBMS. The Committee believes the proposed language is less prescriptive (i.e., more flexibility), yet hopefully still ensures the quality of the analytical data.

In making these proposed changes the Committee has attempted to balance the need for more flexibility in the standards with the desire to not go to far and introduce excessive flexibility that could prove to be too vague or ill-advised. The Committee is currently discussing and considering its proposed language and public comments on the proposed language changes. The Committee is committed to assuring that the NELAC Quality Systems standards provide a foundation for PBMS implementation.

7. Question: Several States have indicated that it is very desirable that a laboratory already be actively analyzing samples for a particular program and by a method for which they want to be accredited. However, these same states have relayed that this ideal scenario is often not the case, as a laboratory may request accreditation in attempts to expand their scope of analytical services or in order to satisfy contractual requirements. These states ask: How will the QS standards help ensure that laboratories will have sufficient data for an onsite assessment especially given the proposed changes to the MDL section?

Answer: The MDL, section D.1.4, in the 1998 NELAC standards has a requirement that "MDLs" be determined initially (40 CFR Part 136, Appendix B) and be verified yearly by the analysis of at least one clean matrix sample spiked at the current reported MDL. Under the proposed revision to Section D.1.4, "Detection Limits" are to be determined initially and each time there is significant change in the test method or instrument type. The proposed standard still requires "MDL" if required in the mandated test method or applicable regulation. If the MDL is not required a "detection limit" must still be determined. Therefor the new section

D.1.4 requirements should still help assure that performance data will be available for review by inspectors. In addition, laboratories are required to successfully complete two out of three PT samples yearly and this data would be available for review, as per section 5.5.4 and Chapter 2). However, under the current PT requirements this may only include one method of multiple methods employed by a laboratory for a given parameter group, e.g., metals.

Laboratories also must perform an Initial Demonstration of Analytical Capability (5.10.2.1, D.1.3 Method Evaluation and Appendix C). This data would be available for on-site review. Also note that the QS committee plans to expand Appendix C (IDC) procedures prior to NELAC V to make it applicable to methods for which spiking is difficult or impossible, e.g., Total Suspended Solids, which should further ensure that performance data is available for review.

In addition under Section 5.6.2.3.c. of QS, the Laboratory Management must ensure that the training of personnel is kept up-to-date, which includes a analyst certification to perform the most recent version of the test method (the approved method or standard operating procedure) and documentation of continued proficiency by at least one of the following once per year: i. acceptable performance of a blind sample (single blind to the analyst); ii. another initial demonstration of method capability; iii. successful analysis of a blind performance sample on a similar test method using the same technology; iv. at least four consecutive laboratory control samples with acceptable levels of precision and accuracy; vi if i-iv cannot be performed, analysis of authentic samples that have been analyzed by another trained analyst with statistically indistinguishable. These requirements should further help assure performance data is available on-site for review.

ACKNOWLEDGMENT LETTER, REVIEW GUIDELINES, and COMMENTER TEMPLATE Ouglity Systems Committee

Quality Systems Committee September 17, 1999

	Date:	
Dear	:	

On behalf of the Quality Systems Committee, thank you for your comments on the Chapter 5 standards of the National Environmental Laboratory Accreditation Conference (NELAC). The standards are routinely reviewed and updated. Continual improvement of the standards is the focal point of NELAC process. We encourage your continued written input as well as your attendance at the NELAC interim meeting and yearly conference. Also, our committee routinely schedules 1-2 open forum meetings during each calender year.

Our committee requests that all comments be supplied in electronic format (WordPerfect if possible) and that handwritten, hardcopy and the use of color fonts be avoided. Comments are considered by the QS committee on a first come basis. We have placed a template (table) for comments on the NELAC Web page, which we hope will ensure that the processes is efficient. With this process we hope that emphasis can be placed on consideration of the comments so that the available time is not spent in the mechanics of exchanging information (US Mail and re-typing comments). Routinely, each set of comments is assigned a QS leader who will complete the comment table including suggested language for any proposed changes to the NELAC standards. The Leader will guide a discussion of the comments during routine committee meetings. The minutes of the meeting (posted on the web site) will capture the information in the completed table from committee discussions, thoughts/rationale and present the final decisions.

Again, thank you for taking the time and effort to improve the NELAC Quality System standards.

Sincerely,

Joseph Slayton, Chair Quality Systems Committee

QS Approach: Comments Received and QS Response:

- 1. A form letter will be sent to each commentor notifying them of receipt of the comment and of the QS's approach to reviewing comments and associated updates to the standards.
- 2. QS will consider the comments in the order received.
- 3. A QS committee member will be designated as the lead on each set (or up-set) of the comments from each commentor, who will provide written comments and who will lead a discussion with the full committee on any proposed changes to the standards (including providing the proposed standard language).
- 4. Proposed changes to the standards will be captured in the QS meeting minutes which are posted on the NELAC Web page.
- 5. All comments and written responses will be attached to QS meeting minutes.
- 6. <u>No colors</u> to be used in the comments nor in the response. Use double underlines for additions and strike-outs for removal of items.
- 7. All comments are to be provided in WordPerfect or rich text format using the following the following table:

GUIDING PRINCIPLES/REVIEW CRITERIA

The QS Committee established a set of criteria by which to evaluate the requirements specified in Chapter 5. The standards in Chapter 5 should meet the criteria listed below:

Flexible:

Allow laboratories freedom to use their experience and expertise in performing their work and allow for new and novel analytical methods and approaches, (e.g., Performance Based Measurement System [PBMS]). That the standards specify the "What" and avoid were possible the "How To", (e.g., control limits must be developed to determine if a QC check result is acceptable, the standards do not specify how the laboratory is to determine these limits).

Auditable:

Sufficient detail is included so that the accrediting authorities evaluate laboratories consistently and uniformly.

Practical/Essential:

The standards are necessary QA policies and QC procedures and that these standards should not place an unreasonable burden upon laboratories.

Widely Applicable:

International scope- consistent with ISO Guide 25. Represent QA policies, which establish essential QC procedures, that are applicable to environmental laboratories regardless of size and complexity.

Appropriate For The Use of the Data:

Helps ensure that associated environmental data is of known quality and that the quality is adequate for the intended use of the data.

Comment ID #: , Source of Comments (Name): QS Lead on Response (Name):					
Standard Rev. # SECTION#	COMMENTwith Rationale to QS	QS Leader Provided	RATIONAL		
and QS Standard Narrative		Proposed Change	(from QS Leader)		
(To Filled in by Commentor)	(To Be Filled in by Commentor)	(Commentor Leave	(Commentor Leave		
	New Wording for Standard	Blank)	Blank)		
	(To Be Filled in by Commentor)				

RESPONSES TO COMMENTS

Quality Systems Committee September 17, 1999

Section	Text	Comments and Proposed Text	QS Leader Provided Change	Rationale
Chapter 5		•		
5.9.4.2	Sentence 4 - "If more stringent standards or requirements are included in a mandated test method or by regulation, the laboratory shall demonstrate that such requirements are met."	Judgement as to whether requirements in mandated test methods are more stringent, rather than just different, can be arbitrary. Alternative: Where mandated test methods and/or regulations for calibration exist, the laboratory will follow calibration requirements as specified in	No Change	Committee added "If it is not apparent which standard is more stringent, then the requirements of the regulation or mandated test method are to be
5.9.4.2.1 f)	"Results of samples not bracketed by initial calibration standards must be reported as having less certainty,"	This sounds as if calibration curves are required before and after sample analysis, rather than suggesting that target analyte results reported outside of calibration range must be flagged. This is not appropriate for all analyses, for example, in ICP analyses a linear range	No change	Current version states "Results of samples not bracketed by an initial instrument calibration standards (within
5.9.4.2.1 g)	Sentence 2 – "Data associated with an unacceptable initial instrument calibration shall not be reported."	There may be some cases where reporting such data with a flag is appropriate and better than any alternative available. For example, if one compound in a multi-analyte initial calibration is slightly outside of QC limits, and re-extraction and/or re-analysis is not an option due to lack of available sample and/or holding time, the data would be valid for all points except the one compound, which could be considered estimated or rejected.	Change to read, "Data associated with an unacceptable initial	Allows the client to decide if the data are "good enough" for the decision that must be made. Potential reduction of resampling costs.
5.9.4.2.2 b)	"A continuing calibration check must be repeated at the beginning and end of each analytical batch."	Test methods that require this practice to ensure accuracy already contain the requirement. For some test methods, other QC data exist to continuously monitor accuracy (e.g. internal standard response, signal to noise ratio, recovery standards, internal standards, surrogate standards, mass spectral data). In addition, where a calibration curve is already	No change	Only required when initial calibration is not performed on the day of analysis.

5.9.4.2.2 c)	"In each analytical batch the calibration verification checks must include concentrations at the lowest and highest concentration of the initial instrument calibration."	alternating them appropriate? The lowest and highest points are not appropriate for monitoring quantitative accuracy. Calculated results are likely to be either slightly below or slightly above the calibration range of the curve. In both cases, the calculated values would,	QS Committee consensus was to make no change Change to read, "In each analytical batch the calibration verification checks must include two (2) different concentrations, one above and one below the midpoint of the initial instrument calibration	The intent of this section is to encourage verification of the initial instrument calibration over the entire calibration range.
5.10.5 b)	"Original containers (such as provided by the manufacturer or vendor) shall be labeled with an expiration date."	This appears to include all containers, such as empty sample containers, rather than just containers of standards and reagents.	No change	Section title specifies "Standards and Reagents".
5.13 17)	"clear identification of numerical results with values below 3.18 times the MDL"	Alternative: "above or below the quantitation limit".	No change	Currect version reads "clear identification of numerical results with

Section	Text	Comments and Proposed Text	QS Leader Provided Change	Rationale QS Leader
B Glossary	Text Legal Chain of Custody Definition.	Because both "Chain of Custody" and "Legal Chain of Custody" are included, there is an inference that one is legal and one is not. Recommend removing legal chain of custody.	Drop the reference to "legal" in chain of custody throughout the standard. See references: Table of contents 5.12.4 Legal/or Evidentiary Chain of Custody	QS Leader The two levels of sample handling are sample custody or tracking and CoC
			The legal-chain of custody records shall establish an intact, f) Legal-chain of custody shall begin at the point established by the federal or state oversight program. 5.12.4.3 Controlled Access to Samples Access to all legal chain of custody samples and subsamples shall be controlled and documented. 5.12.4.4 Transfer of Samples to Another Party Transfer of samples, subsamples, digestates or extracts to another party are subject to all of the requirements for legal chain of custody. Glossary App B	
			Legal Chain of Custody (COC): an unbroken trail of accountability that ensures the physical security of samples, data and records	
B Glossary	Quantitation Limits Definition: "with the confidence level required by the data user."	Alternative: "at a stated degree of confidence."		Changed in a later revision
B Glossary	Matrix: "the component or substrate which contains the analyte of interest."	The substrate may not contain the analyte of interest. Component, also, has connotations of analyte rather than matrix.	No change.	Changed in a later revision

		Alternative: The substrate of the test	
		sample.	
C, IDC, Footnote (1); also D, PBMS Footnote (1)	"Accurate: Based on good laboratory practices consistent with sound scientific principles/practices."	"Good Laboratory Practices" is used as the title of two distinct parts of the CFR, Title 40 Part 160 and Title 40 Part 792. "Based on good laboratory practices" can mean that specific QC requirements have been met under two EPA programs (FIFRA and TSCA). In the context of the sentence, "the data associated with the initial demonstration capability are true, accurate," the term representative could be substituted.	Used in the foot note, the intent is to require a recognized process, an industry standard of practice, etc that in general is considered "good practice without any reference to a regulatory requirement.
D, D.1.1 b2)	"Matrix Spikes: Shall be performed at a frequency of one in 20 samples"	The frequency of matrix Specific QC samples should not, and often cannot (due to lack of sample), be determined by a laboratory. Matrix specific QC sample submission and frequency determination needs to be determined on a site-specific basis and followed by the laboratory client. Alternative: The laboratory shall have procedures for measuring and reporting the effect of the matrix on the method performed.	In this section matrix spike is an alternative to LCS as a positive control. If used in place of an LCS the matrix spike must meet the criteria of an LCS, including frequency.

	selected sample(s) shall be rotated among client samples so that various matrix problems may be	based on lab batches results in the information obtained being unusable most of the time. Laboratory batches of 20 samples comprise various clients and sites and matrix QC samples will provide no useable information about the majority of the samples in the batch. Matrix specific QC samples provide crucial information in determination of bias in sample results. This information can only be obtained by using a site-specific approach in matrix specific QC sample assignment. The laboratory is advised in and agency memo, (Clarification Regarding Use of SW-846 Methods, 8-17-98, OSW), "The Agency further recommends that data users should be routinely provided with the MS/MSD results from only those QC samples associated with the field samples from the same site." Controlling matrix QC frequency based on lab batches can result in underestimation of risk at a specific site, where a data user could assume there is no bias of sample results for samples where there is a significant low bias. Alternative: A laboratory shall have procedures to track client sampling batch and to assist clients in assigning matrix specific QC samples at a frequency of 1 per 20. If clients do not provide MS/MSD at the appropriate	This describes a workable process used to gather information without placing an undue burden on the laboratory.
		frequency, the laboratory will note this in the project report.	
D.1.4 a)	"An MDL"	Alternative: "A Detection limit"	Adds to consistency of the standard.

Various	Inclusion of Field activities	Field activities have proceeded with limited	No change	The field sampling issue
Various		oversight. A field accreditation standard is		will be addressed after
	QC procedures	essential in providing control of all facets of		field sampling becomes
	QO procedures	environmental data generation. While some		an "official" part of
		laboratories provide field sampling		NELAC.
		services, it is appropriate that they be		1122101
		covered by a separate field accreditation		
		standard just as a field services company		
		would be.		
		In some cases, field activities and field QC		
		samples have been addressed in NELAC,		
		putting the laboratory in the difficult		
		situation of policing its clients. The		
		invested parties appropriate to comment on		
		field activity QC criteria have not been part		
		of the NELAC process. Inclusion of field		
		activities into this standard could lead to a		
		field unit attached to a lab holding an		
		accreditation that field services companies		
		could not obtain on their own. Conversely,		
		it could put an additional burden on field		
		units associated with a laboratory that field		
		services companies do not need to meet.		
		Field testing is an activity that needs to be		
		defined, and a decision should be made as		
		to whether this document covers field		
		testing, including that provided by labs,		
		field service companies, and industry.		

	Quality Systems Committee	Source	of Comments: ELAB
	QS Lead on Response: Sh	eila Meyers	April 6, 1999
Standard Rev. # SECTION# Standard Narrative	COMMENTwith Rationale to QS	QS Leader Provided Proposed Change	RATIONAL
10.1 5.9.4.2.	Change "more stringent" to "different". Otherwise NELAC will be superseding existing laws, a situation that will never hold up in court. New Wording for Standard: (see last sentence) If more stringent different standards or requirements	No change.	The goal of NELAC and the standards for Quality Systems 5 is to develop a MINIMUM set of consencus standards that must be met by the regulated community and participating programs. More stringent requirements would be those that clearly demonstrate greater proficiency defining test results and a greater degree of confidence. These would not supersede existing requirements, they would be add ons to minimum procedures. If it cannot be determined which program is more stringent (NELAC or method and'or program), then the laboratory will use the method or regulatory requirement. If they are so different as to not be comparable, then both would be a requirement. An example of NELAC being more stringent would be the requirement of PE samples for drinking water. The drinking water program requires one per year; NELAC requires 2 sucessful completions per year.

	Quality Systems Committee	Source	of Comments: ELAB
QS Lead on Response: Sheila Meye			April 6, 1999
Standard Rev. # SECTION# Standard Narrative	COMMENTwith Rationale to QS	QS Leader Provided Proposed Change	RATIONAL
10.1 5.9.4.2.1 (f)	For certain techniques (i.e., ICP), existing methods allow for a quarterly linear range check and the single point standard does not bracket all of the sample results. This section as written would mean that perhaps 50% of current ICP data would be qualified. New Wording for Standard: Results of samples not bracketed by initial calibration standards, or by quarterly linear range checks in the case of ICP, must be reported as having	No change.	Consensus drives a minimum of 2 calibration concentration standards for quantiation. Data reported out (unqualified) must be withen this high and low standard.
10.1 5.9.4.2.1 (h)	Frequently in risk assessments, the regulatory/decision level is the MDL. It is not analytically feasible to include a standard at or below the MDL in a calibration curve. In general, the requirement for the low standard in the curve should be that it is at the Quantitation Limit. New Wording for Standard: (We recommend adding another sentence.) However, the lowest calibration standard should not be below the Quantitation Limit.	No change	If program or regulatory requirements mandate the use of an MDL however defined, that will exceed the requirements of NELAC standards if more stringent. This is a program requirement issue. Not an issue for NELAP. If decisions will be made concerning a certain concentration level, then it is required that a laboratory be able to "see" to that level in order to make decisions (decision level). Chapter 5 does not us the term MDL or "minimum detection limit". Please refer to definition of "Detection Limit."

	Quality Systems Committee	Source	of Comments: ELAB
	QS Lead on Response: Sheila Meyers		April 6, 1999
Standard Rev. # SECTION# Standard Narrative	COMMENTwith Rationale to QS	QS Leader Provided Proposed Change	RATIONAL
10.1 5.9.4.2.2 ©	As stated this requirement is a significant deviation from the current practice of running one of the middle points in the curve (typically the second or third point). Since many analytical samples tend to fall in the bottom third of the concentration range, the high standard is of less value for a continuing calibration check than those with lower concentrations. Therefore, our recommendation is that the wording be changed. New Wording for Standard:must include two concentrations from the	Change has already been made	ě .

	Quality Systems Committee	Source	of Comments: ELAB
	QS Lead on Response: Sh	eila Meyers	April 6, 1999
Standard Rev. # SECTION# Standard Narrative	COMMENTwith Rationale to QS	QS Leader Provided Proposed Change	RATIONAL
10.1 5.9.4.2.2 (f)	The statement that a second consecutive calibration check must meet criteria is too restrictive. A typical scenario for volatiles is that when a check fails, the existing standard is rerun. If it's still outside acceptance limits, the standard is reprepared from concentrated stock solutions. If this new check standard passes, the analysis is continued in that the problem was due to instability or volatility of the standards and not due to instrument issues. Thus, three checks could be run before running a new calibration. This is particularly necessary for the "gases" that are part of the volatiles analytes. New Wording for Standard: (second sentence)within acceptance criteria, a new check standard may be prepared from a fresh stock standard. If this new standard fails, a new initial instrument calibration must be performed.	No change	The typical scenario is acceptable. The second calibration verification check can be at the end of the batch run. This check can also be used to satisfy the calibration verification check for the next batch. Also there has been a rewrite that states only one is required if internal stardards are used.

	Quality Systems Committee	Source of	of Comments: ELAB
	QS Lead on Response: Sh	eila Meyers	April 6, 1999
Standard Rev. # SECTION# Standard Narrative	COMMENTwith Rationale to QS	QS Leader Provided Proposed Change	RATIONAL
10.1 Page 5B-1 - Batch Definition	This definition includes <u>preparation</u> and <u>analytical</u> batches but does not address tests that do not require sample preparation such as water samples for TOX or volatiles analyses. In such a continuous	No change	Already addressed.
	Process with no sample preparation, the batch should be limited to 20 samples with no 24 hour time limit. Otherwise one many never be able to run a full batch of 20 samples even using a continuous autosampler controlled process, if the run time per sample is too long.		
	New Wording for Standard: (add new paragraph) For analytes that do not require sample preparation such as total organic halogens or volatiles, a batch is composed of 20 samples but the time may exceed 24 hours.		

Charles Dyer
Program Manager
State of New Hampshire
Department of Environmental Services
6 Hazen Drive
P.O. Box 95
Concord, NH 03302-0095

Dear Mr. Dyer:

On behalf of the QS committee I would like to thank you for your letter and the comments from Russell D. Foster, Technical Director, RLI Resource Laboratories, Inc and from SCITEST Laboratory Services (Joann). We request that in future submissions that you employ the comments template that QS's has routinely included with our meeting minutes on the NELAC Web page.

- 1. Definition of Preparation Batch, appendix B, page 5B-1. The QS committee agreed upon 20 samples per batch as being consistent with EPA and good laboratory practices. The batch size, drives the analysis of additional QC samples, e.g., method blank and laboratory control samples. In addition, we too wrestled with the need for a time limit in this criteria. The consensus reached: "...with a maximum time between the start of processing of the first and last sample in the batch to be 24 hours". The stress here is "start of process" and we realize that as manufactures provide various automated (sequential) devices this may be problematic, however additional QC under such an automated scenario should have decreased impact on laboratory throughput.
- 2 Request for clarification of the NELAC standard regarding labeling sample containers (NELAC 5.11.1.a). The letter from the Vermont laboratory indicates "...each of the four (sample) bottles has a distinct label, with two distinct Work Order #3". NELAC (5.11.1.a) requires that: "The laboratory shall assign a unique identification (ID) code to each sample container received in the laboratory. The use of container shape, size or other physical characteristic, such as amber glass, or purple top, is not an acceptable means to identifying the sample". The "distinct label" you have indicated should meet the "unique sample (ID) code" requirement, as long as, "This laboratory code shall maintain an unequivocal link with the unique field ID code assigned each container (5.11.1.b) and "The laboratory ID code shall be entered into the laboratory records and shall be the link that associates the sample with related laboratory activities such as sample preparation or calibration "(5.11.1.d).

Sincerely,

Joseph Slayton, Chair QS Committee

Comment ID #:	Source of Comments: Dianne Terry, Terry Affiliates, LLC	QS Lead on Response (Name): Donivan Porterfield	
Standard Rev. #: Draft 1/13/99 Section # and QS Standard Narrative (Commentor)	COMMENT to QS (commentor)	QS Leader Provided Proposed Change	RATIONALE (from QS Leader)
5.9.3c)/5.9.4.1e) "Glass microliter syringes are to be considered in the same manner as Class A glassware, but must come with a certificate attesting to established accuracy or the accuracy must be initially demonstrated and documented by the lab"	Statement appears following 5.9.3c) and following 5.9.4.1e). Is it really supposed to be in both places? Seems out of place in 5.9.3.	No change.	In current revision 11 (29 April 1999) indicated duplication has been eliminated. Indicated content is now only present in 5.9.4.1.e.
5.12.2.d) "The laboratory shall establish a record management system for control of laboratory notebooks; instrument logbooks; standards logbooks; and records for data reduction, validation storage and reporting."	Re: "validation storage" Is this requirement for a records management system for records for validation and for storage, or for validation storage (whatever that is)? I.e., is there a comma missing? "validation, storage" or possibly it should read "and for storage of records for data reduction, validation, and reporting;"	" validation, storage"	Yes, believe that a comma is missing. The current phrase going back to about version 4.
5.13.a)17) "clear identification of numerical results with values outside quantitation limits.	Every result, unless equal to the quant limit, is outside the quant limit. Does this item mean "values less than the quantitation limit" or is it meant to address results outside the calibration range as in 5.9.4.2.1.f)	No change.	The current wording is to allow for analytical methods that may have both a lower and upper quantitation level, e.g. the upper quantitation level representing the linear response limit of the technquie.
5.9.4.2.1.f) "Results of samples not bracketed by initial calibration standards must be reported as having a less certainty, e.g., defined qualifiers or flags or explained in the case narrative."	What about single point calibration? Or when no calibration is required (re: no calibration for Standard Methods 4500-NH3 E for Ammonia-Ion Selected Electrode Using Known Addition, which is approved for NPDES wastewater analysis) Are these non-multipoint calibrations considered a semiquantitative or qualitative analysis that will by definition require a qualifier since there is no multi-point ICAL. For analysis such as pH, if calibration is checked at pH 10 and the sample is pH 11 must it be qualified? What about titration methods such as for chloride using standardized silver or mercuric nitrate with no multi-point calibration (Standard Methods 4500-Cl B & C)? These should not require	No change.	The proposed version of chapter 5 requires at least two calibration points (5.9.4.2.1.i) for those techniques utilizing an initial calibration. Thus the issue of single point calibration is effectively addressed. Where no calibration is required as in the referenced Standard Method the requirement is moot. For the pH example provided the pH 11 result would need to be reported as having less certainty since an initial calibration is being performed and the example

Comment ID #:	Source of Comments: Dianne Terry, Terry	QS Lead on Response (Name):	
	Affiliates, LLC	Donivan Porterfield	
	reporting with qualifiers to satisfy		result is outside the example
	5.9.4.2.1.f). Suggest clarification for this		calibration range.
	section, such as "Where multi-point initial		
	calibration is required, results of samples		
	not bracketed by initial calibration		
	standards"		
Chapter 1 Policy and Structure, Figure 1-3 =	Is it going to be program/METHOD/analyte	No change.	This topic is not within the
fields of testing, program, method, analyte;	or program/MATRIX/analyte?		jurisdiction of this committee.
Chapter 2, Proficiency Testing 2.1.3 a)b)c)			That question is better directed to
indicates PT fields of testing are program,	Do not propose to change this section		the relevant committee.
matrix type, analyte; and Chapter 5, QS	because I agree that reference to the Chapter		
5.4.2j) refers to Chapter 2: "when available,	2 makes most sense, so you do not end up		
participate in inter-laboratory comparisons	with inconsistencies if one Chapter changes		
and proficiency testing programs. For	and the other does not. But could you get		
purposes of qualifying for and maintaining	some resolution of method vs matrix? Or a		
accreditation, each laboratory shall participate	confirmation that accreditation is per		
in a proficiency test program as outlined in	method, but for purpose of PTs will only		
Chapter 2.0."	need matrix and will apply to more than one		
	method? Would this mean		
	SDWA/WATER/VOA could be analyzed by		
	either GC or GC/MS or is it also meant to be		
	'technology' based (it does not say that)?		
Initial demonstration of method capability:	Could this be called the same thing	No change.	The current revision seeks this
5.6.2.c)3)ii) "initial demonstration of	throughout? Since it is "one of" the		consistency in using the
method capability" (changed, was method	acceptable methods of establishing not only		terminology "Demonstration of
performance); 5.10.2.1 title "Method	initial but ongoing method validation it		Capability" (DOC). This
Validation/Initial Demonstration of	would make more sense to label it as such.		terminology covering both the
Capability"; 5.10.2.1.a) and d) "initial	If you want to stick with "initial		initial and continuing aspects.
demonstration of method performance";	demonstration of capability (IDC)" to be		
Appendix C Initial Demonstration of	consistent with the newer EPA methods,		
Capability, C1 Initial Demonstration of	then use that but label it as a method to		
Capability, C1 first paragraph "initial	achieve the end. Example, as with VOA by		
demonstration of method performance"; C2	GC or VOA by GC/MS, it could indicate		
first paragraph and 'certificate' title = initial	"method validation by IDC" or "method		
demonstration of capability.	performance by IDC" or "demonstration of		
	method capability by IDC" – whatever. See		
	associated comments for Appendix C		
	certificate (next item)		

Comment ID #:	Source of Comments: Dianne Terry, Terry Affiliates, LLC	QS Lead on Response (Name): Donivan Porterfield	
Appendix C, Certificate Statement 6. Title "Initial Demonstration of Capability" 7. "Method Number, and Analyte, or Class of Analytes or Measured Parameters" 8. There is no reference to the lab SOP/revision number used to establish method capability 9. under "We, the undersigned", Item 5. "All raw data…retained at the facility" Example revised certificate attached.	 Title, change to "Method Validation/Initial Demonstration of Capability" Change to "Prep/Analysis Method Number(s)" to ensure the prep method is identified Add a line for Lab SOP No(s)/Rev # Change at the facility to by the facility. These records may be archived by the lab, not necessarily on site, as long as readily retrievable. Or is the argument that the off-site archives are agents of the facility so actually it is retained "at" the facility. 	Appropriate language to address point 3 to be added. Appendix C (Certification Statement), 5: change " retained at the facility" to "retained by the laboratory".	 As noted above the title has been changed. While not specifically calling out "prep" methods the certificate language has been modified to consider the demonstration of multiple methods being certified. Agreed. Agreed.

Comment ID #:	Source of Comments: Dianne Therry, Therry Affiliates, LLC	QS Lead on Response (Name):	
Standard Rev. #: Draft 1/13/99 Section # and QS Standard Narrative (Commentor)	COMMENT to QS (commentor)	QS Leader Provided Proposed Change	RATIONALE (from QS Leader)
5.9.3c)/5.9.4.1e) "Glass microliter syringes are to be considered in the same manner as Class A glassware, but must come with a certificate attesting to established accuracy or the accuracy must be initially demonstrated and documented by the lab"	Statement appears following 5.9.3c) and following 5.9.4.1e). Is it really supposed to be in both places? Seems out of place in 5.9.3.		
5.12.2.d) "The laboratory shall establish a record management system for control of laboratory notebooks; instrument logbooks; standards logbooks; and records for data reduction, validation storage and reporting."	Re: "validation storage" Is this requirement for a records management system for records for validation and for storage, or for validation storage (whatever that is)? I.e., is there a comma missing? "validation, storage" or possibly it should read "and for storage of records for data reduction, validation, and reporting;"		
5.13.a)17) "clear identification of numerical results with values outside quantitation limits.	Every result, unless equal to the quant limit, is outside the quant limit. Does this item mean "values less than the quantitation limit" or is it meant to address results outside the calibration range as in 5.9.4.2.1.f)		
5.9.4.2.1.f) "Results of samples not bracketed by initial calibration standards must be reported as having a less certainty, e.g., defined qualifiers or flags or explained in the case narrative."	What about single point calibration? Or when no calibration is required (re: no calibration for Standard Methods 4500-NH3 E for Ammonia-Ion Selected Electrode Using Known Addition, which is approved for NPDES wastewater analysis) Are these non-multipoint calibrations considered a semiquantitative or qualitative analysis that will by definition require a qualifier since there is no multi-point ICAL. For analysis such as pH, if calibration is checked at pH 10 and the sample is pH 11 must it be qualified? What about titration methods such as for chloride using standardized silver or mercuric nitrate with no multi-point calibration (Standard Methods 4500-Cl B & C)? These should not require		

Comment ID #:	Source of Comments: Dianne Therry,	QS Lead on Response (Name):	
	Therry Affiliates, LLC		
	reporting with qualifiers to satisfy		
	5.9.4.2.1.f). Suggest clarification for this		
	section, such as "Where multi-point initial		
	calibration is required, results of samples		
	not bracketed by initial calibration		
	standards"		
Chapter 1 Policy and Structure, Figure 1-3 =	Is it going to be program/METHOD/analyte		
fields of testing, program, method, analyte;	or program/MATRIX/analyte?		
Chapter 2, Proficiency Testing 2.1.3 a)b)c)			
indicates PT fields of testing are program,	Do not propose to change this section		
matrix type, analyte; and Chapter 5, QS	because I agree that reference to the Chapter		
5.4.2j) refers to Chapter 2: "when available,	2 makes most sense, so you do not end up		
participate in inter-laboratory comparisons	with inconsistencies if one Chapter changes		
and proficiency testing programs. For	and the other does not. But could you get		
purposes of qualifying for and maintaining	some resolution of method vs matrix? Or a		
accreditation, each laboratory shall participate	confirmation that accreditation is per		
in a proficiency test program as outlined in	method, but for purpose of PTs will only		
Chapter 2.0."	need matrix and will apply to more than one		
Chapter 2.0.	method? Would this mean		
	SDWA/WATER/VOA could be analyzed by		
	either GC or GC/MS or is it also meant to be		
	'technology' based (it does not say that)?		
Initial demonstration of method capability:	Could this be called the same thing		
5.6.2.c)3)ii) "initial demonstration of	throughout? Since it is "one of" the		
method capability" (changed, was method	acceptable methods of establishing not only		
performance); 5.10.2.1 title "Method	initial but ongoing method validation it		
Validation/Initial Demonstration of	would make more sense to label it as such.		
Capability"; 5.10.2.1.a) and d) "initial	If you want to stick with "initial		
demonstration of method performance";	demonstration of capability (IDC)" to be		
Appendix C Initial Demonstration of	consistent with the newer EPA methods,		
Capability, C1 Initial Demonstration of	then use that but label it as a method to		
Capability, C1 first paragraph "initial	achieve the end. Example, as with VOA by		
demonstration of method performance"; C2	GC or VOA by GC/MS, it could indicate		
first paragraph and 'certificate' title = initial	"method validation by IDC" or "method		
demonstration of capability.	performance by IDC" or "demonstration of		
	method capability by IDC" – whatever. See		
	associated comments for Appendix C		
	certificate (next item)		
Appendix C, Certificate Statement	1. Title, change to "Method	No change	Is clear enough.
5. Title "Initial Demonstration of Capability"	Validation/Initial Demonstration of		

Comment ID #:	Source of Comments: Dianne Therry,	QS Lead on Response (Name):	
Comment ID #.	Therry Affiliates, LLC	Q3 Lead on Response (Name).	
6. "Method Number, and Analyte, or Class of	Capability"		
Analytes or Measured Parameters"	2. Change to "Prep/Analysis Method		
7. There is no reference to the lab	Number(s)" to ensure the prep method is		
SOP/revision number used to establish	identified		
method capability	3. Add a line for Lab SOP No(s)/Rev #		
8. under "We, the undersigned", Item 5. "All	4. Change at the facility to by the facility.		
raw dataretained at the facility"	These records may be archived by the lab,		
E	not necessarily on site, as long as readily		
Example revised certificate attached.	retrievable. Or is the argument that the		
	off-site archives are agents of the facility		
5.60 (20) (4.1 (6.1)	so actually it is retained "at" the facility.	X 1	
5.6.2.c)3)iv) "At least four consecutive	Does this mean 4 in a row, or in the normal	No change	Is clear enough.
laboratory control samples with acceptable	course of analysis the next 4 LCS, which		
levels of precision and accuracy"	could be, for example, over 3-4 days and		
	interspersed with other standards and		
	samples. Suggest adding sentence similar to		
	wording in Appendix C, C1, c) "the 4 LCS		
	can be prepared and analyzed according to		
	the test method either concurrently or over a		
	period of days, interspersed with calibration		
	standards and samples, but must be		
	consecutive LCS data points."		
5.6.2.c)3)v "If i-iv cannot be performed,	Clarification: What is an "authentic	delete "authentic"	agree with comment
analysis of authentic samples that have been	sample"? Is it meant to be the potentially		
analyzed by another trained analyst with	contaminated field or investigative sample		
statistically identical results indistinguishable	submitted for analysis? Or can it be any		
results."	sample from anywhere as long as you have		
	some kind of confirmation that it is positive		
	for the analyte of interest and some		
	experienced analyst got the "same" answer.		
5.9.4 Calibration, paragraph immediately	Consistent format: either use (1) and (2) in	no change	editiorial
preceding 5.9.4.1. "Calibration requirements	the text or use 1) and 2)		
are divided into two parts: (1) requirements			
for analytical support equipment, and 2)			
requirements for instrument calibration. In			
addition, the requirements for instrument			
calibration are divided into initial instrument			
calibration and continuing instrument			
calibration verification."			
	_		

Comment ID #:	Source of Comments: Dianne Therry,	QS Lead on Response (Name):	
	Therry Affiliates, LLC	• , , ,	
5.12.2 b) "All records, including those	1st sentence- if last use was pulling data ~5	Delete "from last use"	agree
specified in 5.12.3 and 5.12.4, shall be retained	years old from storage for a data request		
for a minimum of five years from last use. All	from the client or a lawyer or whoever, that		
information necessary for the historical	the lab must keep it an additional 5 years		
reconstruction of data must be maintained by	from that point, even if the requestor		
the laboratory. Records which are stored only	"finishes" with it? Should at least indicate		
on electronic media must be supported by the	"from last use by the laboratory, unless		
hardware and software necessary for their	superseded by contractual requirements" or		
retrieval."	lab is stuck guessing when the client has		
	finished with their report.		
	In the last sentence, conversely does this		
	mean that if you hardcopy the electronic		
	records the hardcopy is sufficient record?		
	For example, if mass spec raw data such as		
	tunes, calibration, spectra for hits are all		
	hard copied the tapes do not have to be		
	maintained (assuming you have no project		
	contractual requirements)? In order to not		
	have to keep tapes, would this mean that for		
	every 5 point you would have to have spectra		
	for each compound hard copied if you did		
	not want to maintain electronic records?		
5.4.2f) last sentence "The technical director(s)	4.1.1.1a)c)d). can the specified bachelor's	need to bring to the attention of Chapter 4	
shall meet the requirements specified in the	degree be BS or BA? Is the engineering	Committee	
Accreditation Process (see 4.1.1.1)	degree any kind (chemical, electrical,		
	mechanical, etc)?		
	4.1.1.1d), paragraph 2 and its subsections i)		
	through iii): this whole section should		
	probably be new section 4.1.1f)		

Comment ID #:	Source of Comments: Virginia NELAC Work	group	QS Lead on Respons	e: F.Siegelman
Standard Rev. 10.1 And QS Standard Narrative (To Be Filled in by Commentor)	COMMENT with Rationale to QS (To be filled in by Commentor) New Wording for Standard (To Be Filled in by Commentor)	QS Leader Provided Proposed Change (Commentor Leave Blank)		RATIONAL (from QS Leader) (Commentor Leave Blank)
5.4.2.g. "have a quality assurance officer (however named) who has responsibility for the quality system and its implementation. The quality assurance officer shall have direct access to the highest level of management at which decisions are taken on laboratory policy or resources, and to the technical director. Where staffing is limited, the quality assurance officer may also be the technical director or deputy technical director;"	"have a quality assurance officer (however named) who has responsibility for the quality system and its implementation. The quality assurance officer should have direct access to the highest level of management at which decisions are taken on laboratory policy or resources, and to the technical director. However, it is possible that neither the QAO nor TD may have access to highest levels of management at which decisions are taken on laboratory policy or resources." This is an important issue for many organizations. The QAO may report directly to the Technical Director (TD) but neither the QAO not TD may have access to senior management even though policy decisions are made at this level. The standards must acknowledge that the Technical Director is capable of managing the laboratory and may not have input into the policy making process. In many situations, to comply with this requirement, the entire organization would have to be restructured.	No change		The text: "at which decisions are taken on laboratory policy" defines the level of management that both the QAO and TD need access to which is not necessarily the senior management.
5.4.2.j. "When available Chapter 2."	Clarification is needed. If the PT programs stated is the same as in Ch.2, then this statement is unnecessary since PT is required in Ch.2. Please delete to minimize redundancy. If not, please clarify.	No change		the text: "when available, participate in inter-laboratory comparisons and proficiency testing programs. For purposes of qualifying for and maintaining accreditation, each laboratory shall participate in a proficiency test program as outlined in Chapter 2.0." deals with both PT and inter-laboratory comparisons and references Chapter 2.0 Additionally since PT sample supply is going through a

			transition that includes privatization and since PT samples for all matricies are currently not available, the standard as written addresses the current situation. The deletion would require labs to participate in proficiency testing programs that currently do not exsist.
5.5.1.c. "The laboratory shall define and document its policies and objectives for, and its commitment to accepted laboratory practices and quality of testing services."	This section seems to be saying the same thing as 5.5.2. Would it be more appropriate for this statement to be made in the Quality Manual, which outlines the laboratories policies, procedures and other quality statements? How many places do the statements of commitment to quality need to be confirmed?	No change	I believe we addressesed a comparision of 5.5.1.c and 5.5.2 in the first set of Virginia comments and I concur with the response given then: "5.5.1c The commentor claim that there repetitiveness with 5.2 in the construction of the quality manual. However, there was a misreading. The management statement of objectives is different from that stated for the laboratory Policies and objectives. Indeed, it may have some similarities, but it is different. I am not sure that there is any way to make the contents of these statements clearer. I would retain the wording as it is."
5.5.1.e. "The quality manual shall be"	"The quality assurance officer shall ensure that the quality manual is current."	No Change	Current text adequately addresses need.
	Change from passive to active voice. This QAO responsibility should be moved to the list of QAO responsibilities in section 5.4.2.g.		
5.5.2.a. "a quality statement by top management"	Consolidate quality statements into one document – the Quality Manual. How many quality statements are necessary?	No Change	The current version of the standard is so written. The quality statement is part of the Quality Manual: "The quality

			manual and related quality documentation shall also contain: a) a quality policy statement, including objectives and commitments, by top management;"
5.5.2.c. "the relationship between management, technical operations, support services and the quality system;"	"the relationship between management, technical operations, support services and the quality system e.g. an organizational chart;" Agree with response to previous comments to aid with clarity.	No Change	The inclusion of an organizational chart is already covered elsewhere in the standard.
5.5.2.e. "Job descriptions of key staff and reference to the job descriptions of other staff;"	"job descriptions of key staff and reference to the file location of the job descriptions of support staff" Agree with response to previous comments to aid with clarity.	No Change	The reference can include the file location of other staff but the should not be limited to just that.
5.5.2.j. "reference to the calibration and/or verification test procedures used;"	Please clarify to what calibration and verification test procedures are referring.	No change	Requirements are described in sufficient detail in other parts of the standard and repeating requirements in a document can be confusing to the reader.
5.5.2.n. "reference to verification practices"	Please clarify to what verification practices are referring.	No change	Requirements are described in sufficient detail in other parts of the standard and repeating requirements in a document can be confusing to the reader.
5.5.3.4 Entire section.	Delete section. This is redundant language already described in other chapters. Repeating requirements in a document is confusing to the reader.	No change	While this section does reference other portions of the standard (5.5.4 and chapter 2) this section contains information not found in other places in the standard. It should also be noted that what is offered here is examples and a laboratory's efforts in carrying out Performance Audits does not need to be limited to these.
5.5.4 "are further described in Appendix D."	"Essential quality control procedures shall apply, where applicable, to all testing laboratories. These principles	No change	I concur with the response to a similar comment in the earlier

	and the manner in which they are implemented is dependent on the types of tests performed by the laboratory (e.g) are and further described in Appendix D. Make the first section of Appendix D "General QC Requirements Applicable to all Methods." By consolidating these elements in one place, the reader will not be forced to go back and forth between two sections in the document. This may also prevent misunderstandings and confusion.		Virginia comments: 5.5.4. "are further described in Appendix D." Delete section and move essential QC procedures to Appendix D. The QC procedures only need to be described once, either here or in the appendices. Multiple entries for the same item can result in contradictory statements.
			Response: The intent of Section 5.5.4 is to include the essential QC elements that are applicable to any methods performed under NELAC accreditation regardless of which category of testing is being accredited. Additional elements unique to each category (or not applicable to all categories) of testing are then listed in appendix D. This hierarchial approach is intended to distinguish common elements which cross all categories from those elements specific to a category. Any redundancies listed in appendix D could be deleted, unless it is intended to add clarity to the appendix D section.
5.6.2.c.3. (analyst demonstration of continued proficiency)	Add: "vi. For analyses extending over a period exceeding five days (e.g. WET and bacteriological tests), in which multiple analysts are routinely involved in the analysis of a given sample, analysts shall perform a portion of the analyses described in i-v which is representative of their normal laboratory duties."	The current proposed version of the standard addresses this comment.	This has been addressed in the current version of the standard with changes to the text and the utilization of the concept of a work cell .

Attachment G

	For an analyst to perform an entire chronic WET test, from start to finish including dry weights requires that they work 9 consecutive days. Similarly, the "complete" Coliform test by the fermentation method requires 7 calendar days. Several analysts routinely care for these tests over the entire test period. For example, two or three technicians should be able to conduct a chronic mysid test on a blind sample to satisfy this requirement. Of course if the test fails to meet acceptability requirements, all technicians would be affected.		
5.6.2.d. "Documenting all analytical and operational activities of the laboratory."	"Laboratory management shall be responsible for documenting all analytical and operational activities to demonstrate compliance with the quality system requirements." As written, several interpretations could be made as to what these activities consist of.	No Change	The phrase "Laboratory management shall be responsible for" would be redundant because this section of the standard is titled: "5.6.2 Laboratory Management Responsibilities "The proposed text of "to demonstrate compliance with the quality system requirements." could be interpreted as putting limitations on the management documentation responsibilities.

Policy on Revision of Standards

Once the standards are voted in by the NELAC members, the states and federal agencies (accrediting authorities) have the option of adopting the standards into their statutes, regulations, etc. The accrediting authorities must have some assurance that the standards will have a certain stability over time, in order to successfully implement them within their own jurisdiction. During the period of implementation, especially in the initial phase, there will likely be some issues that are untenable, duplicative, inordinately expensive, etc. It is important that these types of issues be addressed as expeditiously as possible, but that editorial, or other non-essential issues be delayed for a period of time. The following procedures are designed to assure that the standards are revised when appropriate and needed, without constant changes disrupting the adoption and implementation of the standards by the accrediting authorities:

- 1. Each committee chair should review those portions of his or her committee's standards which will be proposed for a vote at the NELAC Annual Meeting.
- 2. Any portions of the revised standards, which have been changed during the two voting sessions immediately preceding the Annual Meeting at which the standards will be put to a vote, must be identified. (For example, those portions of a standard accepted at the 1997 Annual Meeting would be eligible for revision at the 2000 Annual Meeting.)
- 3. During that three (3) year period, committee chairs should assure that any proposed changes are not editorial in nature, e.g. changes to the organization of material, terminology, and syntax.
- 4. If a revision is deemed necessary, the chair must provide the Board of Directors, at least 60 days prior to the Annual Meeting, those revised sections which have not been in effect for at least three (3) years. A justification for the substantive changes, detailing the current problem, its urgency, and the manner in which the remedy will resolve the situation, shall also be provided.
- 5. The Board will review the justification and decide whether the revision is of such a nature to demand immediate action at the next Annual Meeting. Voting on any changes deemed to be editorial will be deferred to a later Annual Meeting.
- 6. Committees are allowed and encouraged to consider editorial changes at any time. However, revisions should only be presented for a vote once the three (3) year time frame is met.
- 7. Each committee chair is responsible to convey to the committee members and any attendees at the Annual and Interim Meetings, the voting restrictions outlined in this policy.

	ommenter's Name: filiation:
Со	mmittee Lead on Response (Name):
1.	Standard Rev. # SECTION# TO BE COMPLETED BY THE COMMENTER: E: Current Standard Text F: COMMENT with Rationale G: Proposed Wording Change
	TO BE COMPLETED BY THE COMMITTEE: H: OUTCOME (Including any proposed change) I: RATIONALE
B:	Standard Rev. # SECTION# TO BE COMPLETED BY THE COMMENTER: C: Current Standard Text D: COMMENT with Rationale E: Proposed Wording Change
	TO BE COMPLETED BY THE COMMITTEE: F: OUTCOME (Including any proposed change) G: RATIONALE
C:	Standard Rev. # SECTION# TO BE COMPLETED BY THE COMMENTER:
	D: Current Standard Text E: COMMENT with Rationale F: Proposed Wording Change TO BE COMPLETED BY THE COMMITTEE:
	G: OUTCOME (Including any proposed change) H: RATIONALE

Date:

Comment ID #: